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GenCore version 5.1.9
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OM protein - protein search, using sw model

Run on:

June 23, 2006, 20:44:14; Search time 39 Seconds (without alignments) 14.803 Million cell updates/sec

US-10-648-854-10 32 1 HAVHAV 6 Title: Perfect score: Sequence: BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

283416 segs, 96216763 residues Searched:

283416 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

1: pirl:* 2: pir2:* 3: pir3:* 4: pir4:* PIR 80:* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed and is derived by analysis of the total score distribution.

SUMMARIES

ū	Description	precorrin-3 methyl	probable spermidin	probable adenylate	probable lacl fami	aspartate transami	hypothetical prote				division	lmbW protein - Str	hypothetical prote	Ω	monoamine transpor	monoamine transpor	probable oligopept	tra protein - Stre	polyketide synthas	hypothetical prote		proteinase I (EC 3	cytochrome-c oxida		hypothetical prote		hypothetical prote	Ω	hypothetical prote	catechol 2,3-dioxy
SUMMERIES	e e	431	H95875	D96025	AP0975	846315	S72847	C70512	B70572	H83133	AF3625	S44970	T37139	S38502	S43686	S43685	G71307	G31844	T17421	AC3046	\$29088	AC3351	T11028	T32399	AG3211	T24600	A71354	858090	H83740	DAAL2E
	DB		~	N	~	~	~	N	~	N	N	~	~	N	~	~	~	~	~	N	N	~	~	~	N	~	7	~	~	-
	Query Match Length		263	649	351	418	514	554	117	209	253	318	345	475	511	511	547	621	1774	164	183	214	261	261	263	265	281	310	315	320
de	Query Match	. 0	100.0	100.0	6.96	6.96	6.96	6.96	90.6	90.6	90.6	90.6	90.6	90.6	90.6	90.6	90.6	90.6	9.06	87.5	87.5	87.5	87.5	87.5	87.5	87.5	87.5	87.5	87.5	87.5
	Score	. "	32	32	31	31	31	31	53	29	29	58	29	29	58	59	59	29	29	28	28	28	28	28	28	58	28	28	28	28
	Result No.	-	73	m	4	ហ	9	7	œ	o,	10	11	12					17	18		20	21	22	23	24	25	56	27	28	29

dihydrodipicolinat probable dihydrodi	MDCR15 protein - h G protein-coupled	G protein-coupled G protein-coupled	hypothetical prote	D-serine dehydrata polysaccharide bio	cardiolipin syntha probable qlycosyl	alpha-amylase (EC D-lactate dehydrod		probable mitogen a
AG3131 D98156	S56162 S26667	S42628 S32785	T08345 E71317	B69965 AG2728	AH2333 H97509	JC5132 AG0144	JC7727	TGSST
0 0	~ ~	~ ~	9 0	010	~ ~	~ ~	01 0	7
321	327	374	397	448	480 522	556 571	575	()
87.5	87.5 87.5	87.5	87.5	87.5	87.5	87.5	87.5	٠. / B
78 78	28 28	58 28	28	28 28	28 28	28 28	28	87
30 31	33 33	3.4 3.5	36	. 80 GH	4 4 0 1	4 4 2 6	44	4. C

ALIGNMENTS

probable spermidineputrescine ABC transporter permease protein SMb20282 [imported] - Sinc C; Species: Sinorhizobium meliloti
R; Finan, T.M.; Weidner, S.; Wong, K.; Buhrmester, J.; Chain, P.; Vorholter, F.J.; Hernand Proc. Natl. Acad. Sci. U.S.A., 98, 9889-9884, 2001
A; Fitle: The complete sequence of the 1,683-kb pSymB megaplasmid from the N2-fixing endon A; Reference number: ASS842; MUID:21396508; PMID:11481431
A; Residues: 1-263 «KDR»
A; Residues: 1-263 «KDR»
A; Residues: 1-263 «KDR»
A; Residues: 1-263 «KDR»
A; Residues: UNIRROT: Q92WQ7; UNIPARC: UPI00000CB48B; GB:ALS91985; PIDN:CAC48672.1;
A; Experimental source: strain 1021, megaplasmid pSymB
B; Galibert, F.; Finan, T.M.; Long, S.R.; Puhler, A.; Abola, P.; Ampe, F.; Barloy-Hubler, pela, D.; Chain, P.; Cowle, A.; Davis, R.W.; Dreano, S.; Federspiel, N.A.; Fisher, R.P.;
L.; Hyman, R.W.; Jones, T.
Science 293, 668-672, 2001
A; Authors: Kahn, D.; Kahn, M.L.; Kalman, S.; Keating, D.H.; Kiss, E.; Komp, C.; Lelaure,

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Gaps

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NyAlternate names: aspartate aminotransferase
C;Species: Medicago sativa (alfalfa)
C;Date: 07-May-1995 #sequence_revision 21-Jul-1995 #text_change 09-Jul-2004
C;Accession: 846315; 818616; $18039
R;Gregerson, R.G.; Miller, S.S.; Petrowski, M.; Gantt, J.S.; Vance, C.P.
R;Gregerson, R.G.; Miller, S.S.; Petrowski, M.; Gantt, J.S.; Vance, C.P.
A;Title: Genomic structure, expression and evolution of the alfalfa aspartate aminotran
A;Reference number: 846315; MUID:94325464; PMID:8049365
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Molecule type: DNA
A;Residues: 1-418 4GRES.
A;Cross-references: UNIPROT:P28011; UNIPARC:UPI00001250CD; EMBL:L25334; NID:g413725; PI
A;Cross-references: UNIPROT:P28011; UNIPARC:UPI00001250CD; EMBL:L25334; NID:g413725; PI
A;Votes: the authors translated the codon TAC for residue 301 as Ile and CAT for residue
R;Udvardi, M.K.; Kahn, M.L.
A;Udvardi, M.K.; Kahn, M.L.
A;Title: Isolation and analysis of a cDNA clone that encodes an alfalfa (Medicago sativ
A;Reference number: S18616; MUID:92092970; PMID:1753949
                  A;Molecule type: DNA
A;Residues: 1-351 <PAR>
A;Cross-references: UNIPARC:UP100005A70F;.GB:AL513382; PIDN:CAD03298.1; PID:g16504918;
C;Genetics:
A;Gene: STY4100
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M.Rosidues: 1-514 <5MT->
A;Rosidues: 1-514 <5MT->
A;Cross-references: UNIPROT:Q49779; UNIPARC:UPI000000435D; EMBL:U00017; NID:g466994; PI
C;Superfamily: protessome complex-related protein
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A;Residues: 2,7RE,'5-707,'D',409-418 <UDV>
A;Cross-references: UNIPARC:UPI00016DE4D; EMBL:X61577; NID:g19570; PIDN:CAA43779.1;
A;Experimental source: var. Ladak
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Date: 19-Mar-1997 #sequence_revision 25-Apr-1997 #text_change 09-Jul-2004
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C;Keywords: aminotransferase; phosphoprotein; pyridoxal phosphate
F;264/Binding site: pyridoxal phosphate (Lys) (covalent) #status predicted
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                                                                                                                                                                                                                                            Length 351;
                                                                                                                                                                                                                                                                                                            0; Indels
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R;Smith, D.R.; Robison, K.
R;Smith, D.R.; Robison, K.
R;Decription: Mycobacterium leprae cosmid B2126.
A;Reference number: S72885
A;Accession: S72847
                                                                                                                                                                                                                                        Score 31; DB;
Pred. No. 43;
1; Mismatches
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                                                                                                                                                                                                                                    Query Match
Best Local Similarity 83.3%;
Matches 5; Conservative
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228 HAIHAV 233
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ||:|||
408 HAIHAV 413
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  probable adenylate cyclase (EC 4.6.1.1) [imported] - Sinorhizobium meliloti (strain 1021 C.) Species: Sinorhizobium meliloti C.) Date: 24-Aug-2001 #sequence_revision 24-Aug-2001 #text_change 09-Jul-2004 C.) Accession: D96025 R.) Accession: D96025 R.) Wong, K.; Buhrmester, J.; Chain, P.; Vorholter, F.J.; Hernan Proc. Natl. Acad. Sci. U.S.A. 99, 9899-9894, 2001
A.) Title: The complete sequence of the 1,683-kb pSymB megaplasmid from the N2-fixing endc A. Reference number: A98842; MUID:21396508; PMID:11481431
A.) Accession: D96025
A.) Status: preliminary
A. Molecule type: DNA
A.) Residues: 1-649 - KURN
A.) Residues: 1-649 - KURN
A.) Cross-references: UNIPROT:0927N9; UNIPARC:UPI00000CB893; GB:AL591985; PIDN:CAC49868.1;
A.) Experimental source: strain 1021, megaplasmid pSymB
A.) Experimental source: strain 1021, megaplasmid pSymB
A.) Experimental source: strain 1021, megaplasmid pSymB
A.) Cross-references: UNIPROT:0927N9; UNIPARC:UPI00000CB893; GB:AL591985; PIDN:CAC49868.1;
A.) Hyman, R.W.; Jones, T.
B.; Hyman, R.W.; Jones, T.
B.; Kallbert, F.; Rahn, D.; Rahn, M.L.; Kalman, S.; Keating, D.H.; Kiss, E.; Komp, C.; Lelaure, hebault, P.; Vandembol, M.: Vorholter, F.J.; Weidner, S.; Wells, D.H.; Wong, K.; Yeh, K.
A.; Title: The composite genome of the legume symbiont Sinorhizobium meliloti.
A. Reference number: A96039; MUID:21368234; PMID:11474104
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B. Parkhill, J.; Dougan, G.; James, K.D.; Thomson, N.R.; Pickard, D.; Wain, J.; Churcher, th, T.; Connerton, P.; Cronin, A.; Davis, P.; Davies, R.M.; Dowd, L.; White, N.; Farrar, Nature 413, 48-852, 201,
A. Authors: Parry, C.; Quail, M.; Rutherford, K.; Simmonds, M.; Skelton, J.; Stevens, K.; A. Title: Complete genome sequence of a multiple drug resistant Salmonella enterica serova, A. Accession: AF0975
A. Status: preliminary
                  ×
hebault, P.; Vandenbol, M.; Vorholter, P.J.; Weidner, S.; Wells, D.H.; Wong, K.; Yeh, A;Title: The composite genome of the legume symbiont Sinorhizobium meliloti. A;Reference number: A96039; MUID:21368234; PMID:11474104
A;Contents: annotation
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                                                                                                                                                                                    A,Gene: SMb20282
A,Genome: plasmid
C,Superfamily: spermidine/putrescine transport system permease protein potI
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Best Local Similarity 100.0%; Pred. No. 50;
Matches 6; Conservative 0; Mismatches
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A;Genome: plasmid
C;Keywords: phosphorus-oxygen lyase
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HAVHAV 480
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Best Local Similarity
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NUCLEOTIDE SEQUENCE.
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Best Local Similarity
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                                                                                   June 23, 2006, 20:40:44; Search time 296 Seconds (without alignments) 18.750 Million cell updates/sec
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02769
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GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
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                                                                                                                                                                                                                                                    2849598 segs, 925015592 residues
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Q3A066_PELCD
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Maximum Match 100%
Listing first 45 summaries
                                                       OM protein - protein search, using sw model
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061AB4
03JR57
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Q37FV8
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03WNS8
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2: uniprot_trembl:*
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Maximum DB seq length: 200000000
                                                                                                                                                US-10-648-854-10
32
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Match Length DB
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                                                                                                                                                              Perfect score:
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brachydanio
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uncultured
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                                                        robaculum
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                                                                            legionella
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                            legionella
                                                                                                                                                                                                                                                                                                                            salmonella
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           STRAIN=DSM 13855;
PubMed=16330755; DOI=10.1073/pnas.0509073102;
PubMed=16330755; DOI=10.1073/pnas.0509073102;
Mongodin B.F., Nelson K.E., Daugherty S., Deboy R.T., Wister J.,
Khouri H., Weidman J., Walsh D.A., Papke R.T., Sanchez Perez G.,
Sharma A.K., Nesbo C.L., Macleod D., Bapteste E., Doolittle W.F.,
Charlebois R.L., Legault B., Rodriguez-Valera F.;
"The genome of Salinibacter ruber: Convergence and gene exchange among hyperhalophilic bacteria and archaea.";
Proc. Natl. Acad. Sci. U.S.A. 102:18147-18152(2005).
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254/3 9SPHI PRELIMINARY; PRT; 133 AA.

D Q254/3;
C Q254/3;
T 24-JAN-2006, integrated into UniProtKB/TrEMBL.
T 24-JAN-2006, entry version 1.
T 07-FEB-2006, entry version 2.
E ISGO5 family transposase orfA.
N GREAMES-SRU_G640;
Salinibacter ruber DSM 13855.
C Bacteria; Bacteroidetes; Sphingobacteria; Sphingobacteriales; C Crenctrichaceae; Salinibacter.
N NCBI_TaxID=309807;
N IL]
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24-JAN-2006, sequence version 1.
07-FEB-2006, entry version 2.
15605 family transposase orfA.
ORFNames-SRU 0191;
Salinibacear ruber DSN 11855.
Bacteria, Bacteroidetes; Sphingobacteria; Sphingobacteriales;
Crenctrichaceae; Salinibacter.
                         Q5x891
Q8zw78
                                                                            052y85
0704a1
0709u4
068gw1
08k278
03ana1
03afi6
05pnfi6
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SEQUENCE 132 AA; 15486 MW; 648013678776A794 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   1 Similarity 100.0%; Score 32; DB Similarity 100.0%; Pred. No. 69; 6; Conservative 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             132 AA.
                   05X891_LEGPA

08ZW78_PRAE

0720431_THETE

070441_SYNPX

06SGW1_9BACT

068KZ78_PROT

03ANA1_SYNSC

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Distributed under the Creative Commons Attribution-NoDerivs License
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    STRAIN=BISBS;
US DOB Joint Genome Institute (JGI-ORNL);
Larimer F., Land M.;
"Annotation of the draft genome of Rhodopseudomonas palustris BisBS.";
Submitted (OCT-2005) to the EMBL/Genbank/DDBJ databases
-:- CAUTION: The sequence shown here is derived from an
EMBL/GenBank/DDBJ whole genome shotgun (WGS) entry which is
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STRAIN=BiBB5;
US DDE Joint Genome Institute (JGI-PGF);
Copeland A., Lucas S., Lapidus A., Barry K., Detter J.C., Glavina T.,
Hammon N., Israni S., Pitluck S., Richardson P.;
"Sequencing of the draft genome and assembly of Rhodopseudomonas
palustris BiBB5.";
Submitted (OCT-2005) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                   STRAIN=ATCC 39073;
US DOE Joint Genome Institute;
Copeland A., Lucas S., Lapidus A., Barry K., Detter J.C., Glaremon I., Israni S., Pitluck S., Chertkov O., Saunders E.H., Brettin T., Bruce D., Han C., Tapla R., Gilna P., Schmutz J., Larimer F., Land M., Kyrpides N., Anderson I., Richardson P.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             100.0%; Score 32; DB 2; Length 193; 100.0%; Pred. No. 1.1e+02; ive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobíales;
Bradyrhizobíaceae; Rhodopseudomonas.
                                       Moorella thermoacetica ATCC 39073.
Bacteria; Firmicutes; Clostridia; Thermoanaerobacteriales;
Thermoanaerobacteriaceae; Moorella group; Moorella.
                                                                                                                                                                                                                                                                                                                   "Complete sequence of Moorella thermoacetica ATCC 39073."; Submitted (DEC-2005) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1 protein.
232 AA; 24151 MW; F26428058EB16451 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                               EMBL; CP000232; ABC20380.1; -; Genomic_DNA.
SEQUENCE 193 AA; 20743 MW; 63BF797Ā36848ED6 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                06-DEC-2005, integrated into UniProtKB/TrEMBL.
06-DEC-2005, sequence version 1.
07-FBB-2006, entry version 3.
07-PBD-2006, entry version 3.
08-Names=RPDDRAFT_0342;
Rhodopseudomonas palustris BisBS.
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NUCLEOTIDE SEQUENCE.
  Peptidase C56, PfpI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
                                                                                                          NCBI_TaxID=264732;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        31 HAVHAV 36
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                                                                                                                                                                                                                               Copyrighted by the UniProt Consortium, see http://www.uniprot.org/terms Distributed under the Creative Commons Attribution-NoDerivs License
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               STRAIN-DSM 13855;
PubMed=16310755; DOI=10.1073/pnas.0509073102;
PubMed=16310755; DOI=10.1073/pnas.0509073102;
Mongodin E.F., Nelson K.E., Daugherty S., Deboy R.T., Wister J.,
Khouri H., Weidman J., Walelb D.A., Papke R.T., Sanchez Perez G.,
Sharma A.K., Nesbo C.L., Macleod D., Bapteste E., Doolittle W.F.,
Charlebois R.L., Legault B., Rodriguez-Valera F.;
"The genome of Salinibacter ruber: Convergence and gene exchange among
hyperhalophilic bacteria and archaea.";
Proc. Natl. Acad. Sci. U.S.A. 102:18147-18152(2005).
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Staplaton M., Carlson J., Chavez C., Frise E., George R., Pacleb J.,
Park S., Wan K., Yu C., Rubin G.M., Celniker S.;
Submitted (OCT-2004) to the EMBL/GenBank/DDBJ databases.
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Bukaryota, Metazoa, Arthropoda, Hexapoda, Insecta, Pterygota, Neoptera, Endopterygota, Diptera, Brachycera, Muscomorpha, Ephydroidea, Drosophilidae, Drosophila.
NCBI_TAXID=7227;
                                                                                                                                                                                                                                                                                                                                                           100.0%; Score 32; DB 2; Length 133; 100.0%; Pred. No. 70;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        100.0%; Score 32; DB 2; Length 138; 100.0%; Pred. No. 73;
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GO; GO:0042302; F:structural constituent of cuticle; IEA
                                                                                                                                                                                                                                                                                               EMBL; CP000159; ABC44828.1; -; Genomic DNA.
SEQUENCE 133 AA; 15851 MW; 923FBE6B892A8CAC CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                138 AA; 14892 MW; 0E2D1525EB301A71 CRC64;
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07-DEC-2004, sequence version 1.
07-FEB-2006, entry version 8.
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Pfam; PP00379; Chitin bind 4; 1.
PRINTS; PR00947; CUTICLE.
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SEQUENCE 138 AA; 14892 MW;
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SEQUENCE.
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Best Local Similarity
Matches 6; Conserv
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QZRGQ9 MOOTH
Q2RGQ9;
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GenCore version
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June 23, 2006, 20:40:25 ; Search time 200 Seconds (without alignments) 13.716 Million cell updates/sec Run on:

US-10-648-854-10 32 Title: Perfect score:

1 HAVHAV 6 Sequence: BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

2589679 seqs, 457216429 residues Searched:

2589679 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

A Geneseq 8:* 1: geneseqmin' •• Database

geneseqp2003bs:* geneseqp2003as:* geneseqp1980s:* geneseqp1990s:* geneseqp2000s:* geneseqp2001s:* geneseqp2002s:* geneseqp2004s:* geneseqp2005s:* Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

geneseqp2006s:*

10:

SUMMARIES

		عد			SOFERE	
Result No.	Score	Query	Query Match Length DB	DB	ID	Description
-1	32	100.0	9		AAY09097	Aay09097 A represe
7	32	100.0	269	۵	ADX67048	
m	32	100.0	401	œ	ADT60801	Adt60801 Plant pol
4	32	100.0	410	4	ABB62935	Abb62935 Drosophil
ហ	32	100.0	266	7	AB069053	Abo69053 Pseudomon
9	32	100.0	615	4	AAM79239	Aam79239 Human pro
7	32	100.0	615	^	ADJ69107	Adj69107 Human hea
æ	32	100.0	645	4	AAM80223	Aam80223 Human pro
σ	32	100.0	1422	~	AAR82066	Aar82066 Hepatitis
10	32	100.0	1422	m	AAB09036	Aab09036 Hepatitis
11	31	96.9	132	~	AAR28990	Aar28990 Encoded b
12	31	96.9	136	ø	ABP99306	Abp99306 Orthosomy
13	31	96.9	215	~	AAR28989	Aar28989 Encoded b
14	31	96.9	257	თ	AEB39698	Aeb39698 L. pneumo
15	31	96.9	283	თ	AEB36280	Aeb36280 L. pneumo
16	31	96.9	487	œ	ADT60381	Pla
17	31	96.9	487	ω	ADT60382	
18	30	93.8	488	œ	ADT60384	Plant
13	29	•	199	2	ADF38938	Adf38938 Corynebac
20	. 53	•	252	^	AB079691	Abo79691 Pseudomon
21	29	٠	268	4	AAM48176	Aam48176 Thermus t
22	53	90.6	268	œ	ADJ67926	Adj67926 T. thermo
23	53	90.6	268	æ	ADJ68138	Adj68138 T. thermo

Adk01216 DNA polym	Adj79435 T. thermo	Adj84875 T. thermo	Adm77663 DNA polym	Adm66330 T. thermo	Ado04383 T. thermo		Adel5667 T. thermo	۲.	Adz76734 T. thermo	Ę,	Aea34165 Thermus t				Adl05393 M. catarr	Adm25638 Hyperther		Abo70399 Pseudomon	Aam99278 Vaccine r	_	Adw35129 HLA bindi
ADK01216	ADJ79435	ADJ84875	ADM77663	ADM66330	ADO04383	ADP82460	ADS15667	ADY55158	AD276734	AEA24742	AEA34165	AEB51569	ADY07855	ABU34984	ADL05393	ADM25638	ABU19998	AB070399	AAM99278	ADW33896	ADW35129
œ	80	œ	œ	œ	œ	80	œ	σ	σ	σ	σ	σ	œ	9	œ	7	9	7	4	7	7
268	268	268	268	268	268	268	268	268	268	268	268	268	350	442	447	9/9	917	1051	11	15	15
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9	90	90	8	8	8	8	8	8	8	90	8	6	90.	8	8	90	8	90	87.	87	87
29	29	29	29	29	53	53	29	29	29	29		29	29	29	53	29	53	29	28		28
24	25	56	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

ALIGNMENTS

RESULT 1

AAY09097 standard; peptide; 6 AA (first entry) 07-JUL-1999 AAY09097; AAY09097

Cadherin; modulating agent; cadherin-mediated cell adhesion; cancer; drug delivery; bladder tumour; ovarian tumour; melanoma; cell adhesion; wound healing; neurite outgrowth; demyelinating neurological disease; multiple sclerosis; carcinoma; leukenis; melanoma; angiogenesis; apoptosis; diabetes; rheumatoid arthritis; immune system; pregnancy; vasopermeability; spinal cord injury; synaptic stability. A representative HAV sequence of a cell adhesion modulating agent.

Synthetic.

WO9916791-A2

08-APR-1999.

98WO-CA000902 29-SEP-1998;

97US-00939853. 29-SEP-1997;

Blaschuck OW, Gour BJ;

(ADHE-) ADHEREX INC.

WPI; 1999-263686/22.

Modulating cadherin-mediated cell adhesion useful for treating neurological disease and cancer.

Disclosure; Page 15; 148pp; English.

The invention relates to methods using cadherin modulating agents, particularly peptides containing the sequence HAV or antibodies, for enhancing or inhibiting cadherin-mediated cell adhesion. The methods can be used to treat diseases or other conditions characterized by undesirable cell adhesion or to facilitate drug delivery to a specific tissue or tumour such as bladder tumours, ovarian tumours or melanomas). They can also be used to enhance cell adhesion (e.g. supplement or replace stitches or to facilitate wound healing). They can also be used

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for enhancing and/or directing neurite outgrowth. The methods can also be used for treating demyelinating neurological disease, e.g. multiple sclerosis. The methods can also be used for e.g. enhancing drug delivery, treating cancers (such as carcinomas, leukemia or melanomas), inhibiting angiogenesis, enhancing adhesion of foreign tissue implanted in a mammal, inducing apoptosis in cadherin-expressing cells, e.g. for treating diabetes or rheumatoid arthritis, modulating the immune system, for preventing pregnancy, increasing vasopermeability, treating spinal cord injuries or inhibiting synaptic stability
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          New recombinant DNA construct, useful for improving plant tolerance to cold, heat, drought, herbicides, extreme osmotic conditions, pathogens or pests, for conferring increased resistance to plant disease, or for
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            The invention describes a recombinant DNA construct comprising a polymucleotide consisting of a sequence encoding an amino acid sequence available in electronic form from the US parent office at ftp. seqdata.uspto.gov/sequence.html?DocID:2004034888. The polymucleotide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              plant protectant; plant growth regulant; gene therapy; plant; recombinant DNA construct; phyadical array; plant breeding marker; cold tolerance, heat tolerance; drought tolerance, herbicide tolerance; extreme osmotic condition; pathogen tolerance; pest tolerance; growth rate; cell cycle pathway; disease resistence; galactomannan production; lighin production; plant growth regulator; yield; plant growth; plant development; seed oil; protein yield;
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                                                                                                                                                                                                      100.0%; Score 32; DB 2; Length 6; 100.0%; Pred. No. 2.1e+06; ive 0; Mismatches 0; Indels
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                                                                                                                                                                                                                                                                                                                                                                                           ADX67048 standard; protein; 269 AA.
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05-NOV-2001; 2001US-00985678.
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Matches 6; Conservative
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KOVALIC D K.
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TABASKA J E.
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(ZHOU/)
(KOVA/)
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(TABA/)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    New recombinant DNA constructs useful in the field of biochemistry and genetics, and in particular for producing transgenic plants with improved biological characteristics.
of the invention are also useful in physical arrays of molecules and as plant breeding markers. The recombinant DNA construct is useful for improving plant tolerance to cold, heat, drought, herbicides, extreme osmotic conditions, pathogens or pests, for manipulating growth rate in plant cells by modification of the cell cycle pathway, for conferring increased resistance to plant disease, for producing galactomannan, lignin or plant growth regulators, for increasing the rate of homologous recombination in plants, for improving yield by modification of photosynthesis or carbohydrate, nitrogen or phosphorus use and/or uptake or by providing improved plant growth and development under at least one stress condition or for modifying seed oil or protein yield and/or content. This is the amino acid sequence of a plant full length insert polypeptide that can be used in the recombinant DNA construct of the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 disease resistance; galactomannan production; plant growth regulator; heat tolerance; herbicide tolerance; lignin production; extreme osmotic condition tolerance; pathogens resistance; pest resistance; yield improvement; seed oil yield; seed protein yield.
                                                                                                                                                                                                                                                                                                                                                                                             Gaps
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                                                                                                                                                                                                                                                                                                                                                Length 269;
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100.0%; Pred. No. 1.9e+02;
ive 0; Mismatches 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADT60801 standard; protein; 401 AA.
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28-APR-2003; 2003US-00425115.
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74 HAVHAV 79
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                                                                                                                                                                                                                                                                                                              Sequence 269 AA;
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                                                                                                                                                                                                                                                                          invention.
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US-10-326-671-50
US-10-05-671-50
US-10-671-419-156
US-10-671-419-156
US-10-671-419-156
US-10-671-134-156
US-10-671-134-156
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US-10-746-167-156
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STATE: Washington
COUNTRY: USA
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Sequence 10458, A
Sequence 10459, A
Sequence 231009,
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Sequence 174117,
Sequence 174117,
Sequence 47481, A
Sequence 47481, A
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37891, A
9644, Ap
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361105,
355177,
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Sequence 10, Appl
Sequence 219794,
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/BMC_Celerra_SIDS3/prodata/2/pubpaa/US08_PUBCOMB.pep:*
/BMC_Celerra_SIDS3/prodata/2/pubpaa/US09_PUBCOMB.pep:*
/BMC_Celerra_SIDS3/prodata/2/pubpaa/US10A_PUBCOMB.pep:*
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15.023 Million cell updates/sec
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                                                                                                                              June 23, 2006, 21:01:15 ; Search time 185 Seconds
                   GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
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US-08-424-550B-81
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                                                                                                                                                                                                                                                                                                                                         2097797 segs, 463214858 residues
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Maximum Match 100%
Listing first 45 summaries
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Gapop 10.0 , Gapext 0.5
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Perfect score:
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Sequence
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                                                                                                                                                                                                   Sequence 10, Application US/09778026
Publication No. US20030013655A1
GENERAL INFORMATION:
APPLICANT: Blaschuk, Orest W.
Gour, Barbara J.
TITLE OF INVENTION: CELL ADHESION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           100.0%; Score 32; DB 3; Length 6; 100.0%; Pred. No. 1.9e+06; ive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/778,026
FILING DATE: 05-Feb-2001
CLASSIFICATION: <Unknown>
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                  NUMBER OF SEQUENCES: 29
CORRESPONDENCE ADDRESS:
ANDRESSEE: SEED and BERRY LLP
STREET: 6300 Columbia Center, 701 Fifth Avenue
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     REFERENCE/DOCKET NUMBER: 100086.402 TELECOMMUNICATION INFORMATION:
                                                                                         ALIGNMENTS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TOPOLOGY: linear
SEQUENCE DESCRIPTION: SEQ ID NO: 10:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       NAME: Maki, David J.
REGISTRATION NUMBER: 32,391
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TELEPHONE: (206) 622-4900
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Squence 37891, Application US/10425114

Squence 37891, Application US/10425114

GENERAL INFORMATION:

APPLICANT: Liu, Jingdong

APPLICANT: Zhou, Yihua

APPLICANT: Zhou, Yihua

APPLICANT: Zhou, Yihua

APPLICANT: Cao, Yongwei

APPLICANT: Cao, Yongwei

APPLICANT: Cao, Yongwei

TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

CURRENT APPLICATION NUMBER: US/10/425,114

CURRENT FILING DATE: 2003-04-28

NUMBER OF SEQ ID NOS: 73128

SEQ ID NO 37891

LENGTH: 269
                                                                                                                                                                                                                                                                    Sequence 62121, Application US/10767701
Publication No. US20040172684A1
Sequence 62121, Application No. US20040172684A1
SEMERAL INFORMATION:
APPLICANT: Kovalic, David K.
APPLICANT: Zhou, Yihua
APPLICANT: Cao, Yongwei
TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated With
TITLE OF INVENTION: Plants and Uses Thereof For Plant Improvement
FILE REFERENCE: 38-21(53535)8
CURRENT APPLICANTION NUMBER: US/10/767,701
CURRENT FILING DATE: 2004-01-29
NUMBER OF SEQ ID NOS: 63128
SEQ ID NO 62121
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                                        100.0%; Score 32; DB 4; Length 74; 100.0%; Pred. No. 65;
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US-10-425-114-37891
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                                                                                    0; Mismatches
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US-10-767-701-62121
                    Query Match
Best Local Similarity luv...
Si Conservative
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Matches 6; Conservative
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ORGANISM: Sorghum bicolor
                                                                                                                                                              27 HAVHAV 32
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ORGANISM: Zea mays
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US-10-424-599-219794
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US-10-425-114-37891
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US-10-767-701-62121
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Publication No. US20040031072A1
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: La Rosa Thomas J
APPLICANT: Kovalic David K
APPLICANT: Thou Yihua
APPLICANT: Cao Yongwei
TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
FILE REFERENCE: 38-21(53223)B
CURRENT APPLICATION NUMBER: US/10/424,599
CURRENT FILING DATE: 2003-04-28
NUMBER OF SEQ ID NOS: 285684
SEQ ID NO 219794
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                                                         Sequence 10, Application US/10648854
Publication No. US2004132651A1
GENERAL INFORMATION:
APPLICANT: Blaschuk, Orest W.
Gour, Barbara J.
TILE OF INVENTION: COMPOUNDS AND METHODS FOR REGULATING
                                                                                                                                                                                                                                                                                                                                                              COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS

SOFTWARE: Patentin Release #1.0, Version #1.30

CURRENT APPLICATION DATA:

FILING DATE: 25-Aug-2003

CLASSIFICATION: <UNknown>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 6;
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OTHER INFORMATION: Clone ID: PAT_MRT3847_40501C.1.pep
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            100.0%; Score 32; DB 4; I 100.0%; Pred. No. 1.9e+06;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               NAME: Jeffrey E. Hundley
REGISTRATION NUMBER: 42,676
REFERENCE/DOCKET NUMBER: 100086.402C2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (206) 622-4900
TELEPHONE: (206) 682-691
INFORMATION FOR SEQ ID NO: 10:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                          NUMBER OF SEQUENCES: 29
CORRESPONDENCE ADDRESS:
ADDRESSEE: Seed IP Law Group PLLC
STREET: 701 Fifth Avenue, Suite 6300
CITY: Seattle
STATE: Washington
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TYPE: amino acid
STRANDEDNESS: «Unknown»
TOPOLOGY: linear
US-10-648-854-10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ATTORNEY/AGENT INFORMATION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity 100.
                                                                                                                                                                                                                                                                                                                                            COUNTRY: USA
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ORGANISM: Glycine max
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US-10-424-599-219794
                 RESULT 2
US-10-648-854-10
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Best Local Similarity
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RESULT 2
US-10-953-349-26203
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Query Match
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Sequence 26203, A
Sequence 26472, A
Sequence 38148, A
Sequence 30764, A
Sequence 361, App
Sequence 361, App
Sequence 360, App
Sequence 41714, A
Sequence 52930, A
Sequence 39938, A
Sequence 39938, A
Sequence 39938, A
Sequence 39937, A
Sequence 39938, A
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                                                                                                                                                                                                                                                                                                           Published Applications AA New:*

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2: /EMC_Celerra_SIDS3/ptcdata/1/pubpaa/US06_NEW_PUB.pep:*
3: /EWC_Celerra_SIDS3/ptcdata/1/pubpaa/US07_NEW_PUB.pep:*
4: /EMC_Celerra_SIDS3/ptcdata/1/pubpaa/US08_NEW_PUB.pep:*
5: /EWC_Celerra_SIDS3/ptcdata/1/pubpaa/US10_NEW_PUB.pep:*
6: /EWC_Celerra_SIDS3/ptcdata/1/pubpaa/US10_NEW_PUB.pep:*
7: /EWC_Celerra_SIDS3/ptcdata/1/pubpaa/US11_NEW_PUB.pep:*
8: /EWC_Celerra_SIDS3/ptcdata/1/pubpaa/US11_NEW_PUB.pep:*
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6.231 Million cell updates/sec
                                                                     June 23, 2006, 21:01:40 ; Search time 22 Seconds
        GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd
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US-10-953-349-26203
US-10-953-349-26203
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US-10-449-902-28647
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US-11-289-102-230
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US-10-953-349-8499
US-10-953-349-33173
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Maximum Match 100%
Listing first 45 summaries
                                                  OM protein - protein search, using sw model
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Gapop 10.0 , Gapext 0.5
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Maximum DB seq length: 200000000
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Perfect score:
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No.
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Sequence 26203, Application US/10953349
Publication No. US20060107345A1
GENERAL INFORMATION:
GENERAL INFORMATION:
TITLE OF INVENTION: SEQUENCE-DETERMINED DNA FRACMENTS AND CORRESPONDING POLYPEPTIDES
TITLE OF INVENTION: ENCONDED THERBY
TITLE OF INVENTION: US/10/953,349
CURRENT APPLICATION NUMBER: US/10/953,349
CURRENT FILING DATE: 2004-09-30
NUMBER OF SEQ ID NOS: 40252
SOFTWARE PATENTIN Version 3.3
SEQ ID NO 26203
LENGTH: 384
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US-10-953-349-26204
US-10-953-349-26204
Sequence 26204, Application US/10953349
Publication No. US20060107345A1
GENERAL INFORMATION:
TITLE OF INVENTION: SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES
TITLE OF INVENTION: ENCONDED THERBY
TITLE OF INVENTION: ENCONDED THERBY
TITLE OF INVENTION: US/10/953,349
CURRENT APPLICATION UNMERS: 2004-09-30
NUMBER OF SEQ ID NOS: 40252
SOFTWARE: PatentIn version 3.3
SOFTWARE: PatentIn version 3.3
SEQ ID NO 26204
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         Sequence 34847, Sequence 8390, A Sequence 8389, A Sequence 3885, A Sequence 3361, Sequence 3227, Sequence 3227, Sequence 52691, Sequence 40819, Sequence 34846, Sequence 32591, Sequence 32590, Sequence 32590
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83.3%; Pred. No. 25;
live 1; Mismatches 0; Indels
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US-10-953-349-33172
US-10-953-349-34847
US-10-953-349-38050
US-10-953-349-38050
US-10-953-349-8189
US-10-953-349-8189
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US-10-953-349-34846
US-10-953-349-32591
US-10-953-349-32590
US-10-953-349-32599
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ORGANISM: Triticum aestivum
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Sequence 362, Application US/10953349
Publication No. US20060107345A1
APPLICATION NO. US20060107345A1
APPLICANT: ALEXANDROV, Nickolai et al.
TITLE OF INVENTION: SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES
TITLE OF INVENTION: SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES
TITLE OF INVENTION: SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES
CURRENT APPLICATION NUMBER: US/10/953,349
CURRENT FILING DATE: 2004-09-30
NUMBER OF SEQ ID NOS: 40252
SOFTWARE: Patentin version 3.3
SEQ ID NO 362
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           APPLICANT: ALEXANDROV, Nickolai et al.
TITLE OF INVENTION: SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES
TITLE OF INVENTION: SENCENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES
FILE REFERENCE: 2750-15799U32
CURRENT APPLICATION WHORE: US/10/953,349
CURRENT APPLICATION WHORE: 2004-09-30
NUMBER OF SEQ ID NOS: 40252
SOFTWARE: Patentin version 3.3
SEQ ID NO 30764
LENDIH: 281
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                                                             APPLICANT: National Institute of Agrobiological Sciences.
APPLICANT: National Institute of Agrobiological Sciences.
APPLICANT: Bio-oriented Technology Research Advancement Institution.
APPLICANT: The Institute of Physical and Chemical Research.
APPLICANT: Foundation for Advancement of International Science.
ITLE OF INVENTION: FULL-LENGTH PLANT CDNA AND USES THEREOF
FILE REFRERNCE: MOA-ADOS571-US
CURRENT APPLICATION NUMBER: US/10/449,902
CURRENT APPLICATION NUMBER: US/10/449,902
PRIOR APPLICATION NUMBER: US 2002-203269
PRIOR PILING DATE: 2002-05-30
PRIOR PILING DATE: 2002-05-30
PRIOR APPLICATION NUMBER: UP 2002-383870
PRIOR APPLICATION NUMBER: UP 2002-11-11
NUMBER OF SEQ ID NOS: 56791
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Pred. No. 47;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ; Sequence 30764, Application US/10953349; Publication No. US20060107345A1; GENERAL INPORMATION:
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; ORGANISM: Triticum aestivum
US-10-953-349-30764
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Best Local Similarity 83.5.
Then 5; Conservative
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Best Local Similarity luv...
5, Conservative
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US-10-449-902-38148
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    94 HAVHVV 99
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US-10-953-349-30764
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US-10-953-349-362
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                                                                                                                                                                                                                                                                                                                    Sequence 26202, Application US/10953349
Publication No. US20060107345A1
GENERAL NO. US20060107345A1
GENERAL NO. US20060107345A1
GENERAL NO. US20060107345A1
APPLICANT: ALEXANDROV, Nickolai et al.
TITLE OF INVENTION: ENCONED THERBY
FILLE OF INVENTION: ENCONED THERBY
FILLE OF INVENTION: ENCONED THERBY
FILLE REPERENCE: 2750-1579PUS2
CURRENT APPLICATION NUMBER: US/10/953,349
CURRENT FILLING DATE: 2004-09-30
NUMBER OF SEO ID NOS: 40252
SOFTWARE: Patentin version 3.3
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Sequence 28667.

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                                         0; Indels
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Best Local Similarity 83.3%; Pred. No. 41;
Matches 5; Conservative 1; Mismatches
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US-10-449-902-38148
; Sequence 38148, Application US/10449902
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ; TYPE: PRT
; ORGANISM: Triticum aestivum
US-10-953-349-26202
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity 83.3
Matches 5; Conservative
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US-10-449-902-28647
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235 HALHAV 240
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266 HALHAV 271
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                                                                                                   1 HAVHAV 6
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US-10-953-349-26202
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LENGTH: 415
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Sequence 37862, A
Sequence 53079, A
Sequence 17799, A
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Sequence 29952, 1
Sequence 19613, 2
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                                                                                June 23, 2006, 20:49:15; Search time 51 Seconds (without alignments) 10.298 Million cell updates/sec
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/ENC_Celerra_SIDS3/ptodata/2/iaa/6_COMB.pep:*
/ENC_Celerra_SIDS3/ptodata/2/iaa/7_COMB.pep:*
/ENC_Celerra_SIDS3/ptodata/2/iaa/H_COMB.pep:*
/ENC_Celerra_SIDS3/ptodata/2/iaa/PCTUS_COMB.pep:*
/ENC_Celerra_SIDS3/ptodata/2/iaa/RE_COMB.pep:*
/ENC_Celerra_SIDS3/ptodata/2/iaa/RE_COMB.pep:*
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GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
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US-09-270-767-37862
US-09-252-991A-17799
US-09-949-016-7142
US-08-469-260A-81
US-08-469-260A-81
US-08-469-260A-81
US-08-469-260A-81
US-08-469-260A-81
US-09-270-767-35705
US-09-270-767-35705
US-09-270-767-35705
US-09-252-991A-28437
US-09-818-780-94
US-09-116-9648-156
US-09-716-9648-156
US-09-252-991A-19145
US-08-505-377-4
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US-09-252-991A-29952
US-09-252-991A-19613
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Maximum Match 100%
Listing first 45 summaries
                                                        OM protein - protein search, using sw model
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Gapop 10.0 , Gapext 0.5
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Maximum DB seq length: 200000000
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32
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Perfect score:
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Sequence 66, Ap
Sequence 200, A
Sequence 6, App
Sequence 1025,
Sequence 21851,
Sequence 31122,
                   Sequence 6
Sequence 6
Sequence 6
Sequence 6
Sequence 6
Sequence 6
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Sequence
Sequence
US-08-202-056-5
US-08-076-093A-6
US-08-284-586-6
US-08-805-478-6
US-08-805-67A-6
US-08-801-228-6
US-08-801-228-6
US-08-801-228-6
US-08-170-496D-66
US-09-170-496D-66
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Sequence 10, Application US/08939853A;
Fatent No. 6203788;
GENERAL INFORMATION:
APPLICANT: Blaschuk, Orest W.
APPLICANT: Blaschuk, Orest W.
TITLE OF INVENTION: COMPOUNDS AND METHODS FOR REGULATING
TITLE OF INVENTION: CELL ADHESION
NUMBER OF SEQUENCES: 30
CORRESPONDENCE ADDRESS:
ADDRESSEE: SEED and BERRY LLP
STREET: 6300 Columbia Center, 701 Fifth Avenue
CITY: Seattle
STATE: Washington
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         100.0%; Score 32; DB 2; Length 6; 100.0%; Pred. No. 5e+05; ive 0; Mismatches 0; Indels
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ZIP: 98104
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
COMPUTER: IBM PC COMPATIBLE
COMPUTER: DatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/939,853A
FILING DATE: 29-SEP-1997
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Maki, David J.
REGISTRATION NUMBER: 32,391
REPERENCE/DOCKET NUMBER: 32,391
RECOMMUNICATION INFORMATION:
TELEPHONE: (206) 622-4900
TELEFRAX: (206) 622-4900
TELEFRAX: (206) 622-4900
TELEFRAX: (206) 622-4900
SEQUENCE CHARACTERISTICS:
SEQUENCE CHARACTERISTICS:
TENTIFY GAMEN 
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TYPE: amino acid
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Matches 6; Conserv
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1 HAVHAV 6 HAVHAV

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Length 566;

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GENERAL INFORMATION:
APPLICANT: VENTER, J. Craig et al.
TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF FILE REFERENCE: CL001307
CURRENT APPLICATION NUMBER: 60/241,755
PRIOR APPLICATION NUMBER: 60/241,756
PRIOR APPLICATION NUMBER: 60/231,768
PRIOR PLING DATE: 2000-10-03
PRIOR FILING DATE: 2000-0-03
PRIOR FILING DATE: 2000-0-09
PRIOR FILING DATE: 2000-0-09
REALOR APPLICATION NUMBER: 60/231,498
PRIOR FILING DATE: 2000-09-08
NUMBER OF SEQ ID NOS: 207012
                                                                                                                                                                                                                                                                                                                                                                                                                                                     APPLICANT: Burges et al.

TITLE OF INVENTION: No. 6989232el Proteins and Nucleic Acids Encoding Same FILE REFERENCE: 21402-099
FILE REPERENCE: 21402-099
CURRENT APPLICATION NUMBER: US/09/939,853A
CURRENT PILING DATE: 2001-08-27
PRIOR FILING DATE: 2000-08-25
PRIOR FILING DATE: 2001-02-08
PRIOR PLICATION NUMBER: 60/267,300
PRIOR FILING DATE: 2001-02-08
PRIOR APPLICATION NUMBER: 60/269,961
PRIOR PILING DATE: 2001-02-20
PRIOR FILING DATE: 2001-03-20
PRIOR FILING DATE: 2001-03-20
                                                                                                    100.0%; Score 32; DB 2; Length 56
100.0%; Pred. No. 1.5e+02;
ive 0; Mismatches 0; Indels
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Patent No. 6812339
                                                                                                                                                                                                                                                                                                                                                                        Sequence 16, Application US/09939853A Patent No. 6989232 GENERAL INFORMATION:
; TYPE: PRT
; ORGANISM: Pseudomonas aeruginosa
US-09-252-991A-17799
                                                                          Query Match
Best Local Similarity 100.v.
-hea 6; Conservative
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SOFTWARE: Patentin Ver. 2.1
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Best Local Similarity 100.
Matches 6; Conservative
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US-09-939-853A-16
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173 HAVHAV 178
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56 HAVHAV 61
                                                                                                                                                                                                         1 HAVHAV 6
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US-09-939-853A-16
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LENGTH: 623
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TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PSEUDOMONAS TITLE OF INVENTION: ARRUGINOSA FOR DIAGNOSTICS AND THERAPEUTICS

TITLE OF INVENTION: ARRUGINOSA FOR DIAGNOSTICS AND THERAPEUTICS

FILE REFERENCE: 10.196-136

CURRENT APPLICATION NUMBER: US 60/074,788

PRIOR APPLICATION NUMBER: US 60/094,190

PRIOR APPLICATION NUMBER: US 60/094,190

PRIOR APPLICATION NUMBER: US 60/094,190

NUMBER OF SEQ ID NOS: 33142

SEQ ID NO 17799

LENGTH: 566
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US-09-270-767-53079
; Sequence 53079, Application US/09270767
; Patent No. 6703491
; GENERAL INFORMATION:
; APPLICANT: Homburger et al.
; TITLE OF INVENTION: Nucleic acids and proteins of Drosophila melanogaster
; FILE REPERRICE: FILE Reference: 7326-094
; CURRENT APPLICATION NUMBER: US/09/270,767
; CURRENT FILING DATE: 1999-03-17
; NUMBER OF SEQ ID NOS: 62517
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 53079
: LENGTH: 107
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                                                                                                                  GENERAL INFORMATION:
APPLICANT: Homburger et al.
APPLICANT: No. Nuclea and proteins of Drosophila melanogaster
FILE REPERENCE: File Reference: 7326-094
CURRENT FPLING DATE: 1999-03-17
NUMBER OF SEQ ID NOS: 62517
SOFTWARE: PatentIn Ver. 2.0
SEQ ID NO 37862
LENGTH: 107
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100.0%; Score 32; DB 2; Length 107;
Best Local Similarity 100.0%; Pred. No. 26;
Matches 6; Conservative 0; Mismatches 0; Indels
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Patent No. 6551795
                                                                       Sequence 37862, Application US/09270767
Patent No. 6703491
                                                                                                                                                                                                                                                                                                                                                                                                        ; ORGANISM: Drosophila melanogaster US-09-270-767-37862
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Best Local Similarity 100.(
Matches 6, Conservative
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81 HAVHAV 86
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